# **BEAD FILLER RUBBER COMPOSITION**

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## Abstract of JP 55054337 (A)

PURPOSE:An extremely hard rubber composition usable for bead filler rubber, having improved rigidity and durability, comprising a (modified) novolak phenolic resin and carbon black. CONSTITUTION:A composion comprising (A) 100 parts by wt. of natural polyisoprene, polybutadiene, or styrene-butadiene copolymer rubber, or their blend, (B) 40-170 parts by wt., preferably 60-120 parts by wt., of carbon black having a dibutyl phthalate absrption &It;=130ml/100g and an iodine adsorption of 40-130mg/g, (C) 15-45 parts by wt., preferably 20-40 parts by wt. based on 100 parts by wt., of (B) of a (modified) novolak phenolic resin, and (D) a curing agent, e.g. hexamethylenetetramine. The use of a movolak phenolic resin and its modified one at a weight ratio of 80:20-20:80 improves the durability synergistically.

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# Miyake et al.

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[54]	BEAD FIL	LER RUBBER COMPOSITION
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[63]	Continuation doned.	n of Ser. No. 85,346, Oct. 16, 1979, aban-
[30]	Foreign	Application Priority Data
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[51] [52]	U.S. Cl	
[58]	Field of Sea	rch 524/495, 511; 525/139; 152/362 R; 260/727, 775, 779 R

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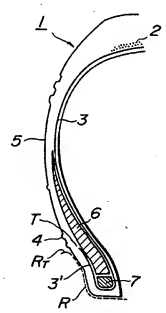
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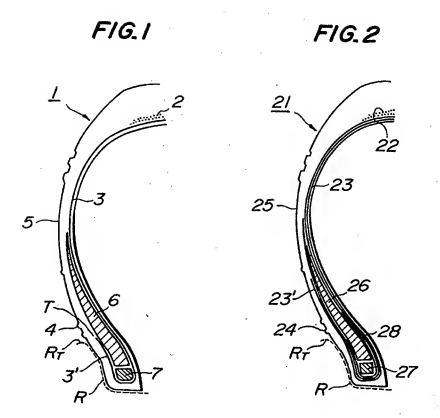
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# [57] ABSTRACT

A rubber composition comprising rubber, novolak-type phenolic resin and carbon black is very rigid and is adapted to be used as a bead filler rubber of a tire. Tires using the rubber composition are excellent in the high speed performance, lateral rigidity, ride feeling and durability.

# 8 Claims, 2 Drawing Figures





## BEAD FILLER RUBBER COMPOSITION

This is a continuation of application Ser. No. 85,346, filed Oct. 16, 1979, now abandoned.

# BACKGROUND OF THE INVENTION

## 1. Field of the Invention

The present invention relates to a rubber composition, and more particularly, relates to a super rigid rub- 10 ber composition comprising rubber, novolak-type phenolic resin and carbon black, and adapted to be used as a bead filler rubber of a tire.

## 2. Description of the Prior Art

There have been variously investigated the structure 15 have accomplished the present invention. of bead portion of radial tire in order to satisfy the rigidity and durability required to tire. For example, it has been attempted to improve the dynamic performance and durability of tire by arranging a bead-reinforcing layer in the bead portion. However, this method 20 has drawbacks that a large number of production steps are necessary and the productivity of the tire is very poor.

Japanese Utility Model Application No. 16,084/72, French Patent No. 1,260,138 and U.S. Pat. No. 25 4,067,373 disclose methods for improving the running performance and other property by arranging super rigid rubber in the bead portion. However, these methods do not substantially think of rubber, which can develop fully a function as a bead filler rubber subjected 30 to complicated forces during the running of a tire and can give a sufficiently high durability to a rubber tire.

While, it is well known to use novolak-type phenolic resin in the production of rigid rubber compositions. However, these rubber compositions substantially con- 35 cern nitrile rubber and neoprene rubber having a high compatibility with the resin. The nitrile rubber series of neoprene rubber series rigid rubber is very difficult to be vulcanized together with natural rubber, polybutadiene rubber and the like, which are commonly used as a 40 rubber for tire, and therefore when the nitrile rubber series or neoprene rubber series rigid rubber is used as a tire part, the rubber is apt to be separated from natural rubber, polybutadiene rubber or the like, and can not be practically used.

In order to solve the above described drawbacks, the inventors have variously investigated how to produce a super rigid rubber composition by compounding novolak-type phenolic resin to natural rubber, polybutadiene rubber or the like, and found out the following facts. 50 Novolak-type phenolic resin is essentially incompatible with natural rubber and other rubbers. Therefore, novolak-type phenolic resin is formed into spherical agglomerates during the kneading commonly carried out in the production of rubber, and has the filling effect only. 55 This phenomenon still occurs even when various resins are used as a resin in place of novolak-type phenolic resin for producing a homogenous mixture of the resin with the rubber. However, when novolak-type phenolic resin is compounded to natural rubber or other rubber 60 together with carbon black, the mixture of the resin and carbon black exhibits a reinforcing effect on the rubber, which is completely different from the reinforcing effect of the resin alone on the rubber or the reinforcing effect of the carbon black alone on the rubber, depend- 65 ing upon the mixing ratio of the resin to the carbon black, and a super rigid rubber composition having a durability remarkably superior to that of conventional

resin-reinforced rubber can be obtained. That is, when natural rubber or diene series rubber is mixed with novolak-type phenolic resin, which is essentially incompatible with the rubber, the resin is separated from the rubber to form island-like large agglomerates in the mixture, but when a mixture of novolak-type phenolic resin and carbon black is compounded to natural rubber or other diene series rubber, the resin disperses uniformly in the rubber without forming large agglomerates similarly to the dispersed state in an ordinary mixture of rubber and carbon black. This action of carbon black to the resin is highly influenced by the mixing ratio of the carbon black to the resin and by the kind of the carbon black. Based on the discovery, the inventors

## SUMMARY OF THE INVENTION

The feature of the present invention is the provision of a bead filler rubber composition, comprising 100 parts by weight of a rubber selected from the group consisting of natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene copolymer rubber and blends thereof, 40-130 parts by weight of carbon black, 15-45 parts by weight based on 100 parts by weight of the carbon black of at least one of novolaktype phenolic resin and novolak-type modified phenolic resin, and a hardener for the resin.

# BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a cross-sectional view of the side portion of a tire according to the present invention; and FIG. 2 is a cross-sectional view of the side portion of a conventional tire.

## DESCRIPTION OF THE PREFERRED **EMBODIMENTS**

The rubber to be used in the present invention includes natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene copolymer rubber and blends thereof. Carbon black is compounded to the rubber in an amount of 40-130 parts by weight, preferably 60-120 parts by weight, more preferably 65-85 parts by weight, based on 100 parts by weight of the rubber. The use of carbon black in an amount of less than 40 parts by weight is too small to disperse homogeneously the necessary amount of resin for reinforcing the rubber. While, when the amount of carbon black is more than 130 parts by weight, the resulting rubber composition is brittle and is very poor in the durability. Further, in the present invention, carbon black having an iodine adsorbability (IA) of 40-130 mg/g and a dibutyl phthalate absorbability (DBP) of not more than 130 ml/100 g defined in ASTM D 1765 is preferably used. Carbon black having an IA of less than 40 mg/g affects adversely the dispersibility of the resin, and carbon black having an IA of more than 130 mg/g is poor in the dispersibility in itself. Therefore, the use of such carbon black is not preferable. When carbon black has a DBP of more then 130 ml/100 g, the carbon black used in an amount sufficient to disperse the resin can not be fully dispersed in the rubber.

In the present invention, the above described carbon black is used together with at least one of novolak-type phenolic resin and novolak-type modified phenolic resin. The novolak-type phenolic resin includes novolak-type phenol resin, novolak-type cresol resin and novolak-type resorcinol resin. The novolak-type modified phenolic resin includes resins obtained by modify-

ing the above described novolak-type phenolic resin with oils, such as rosin oil, tall oil, cashew nut oil, linoleic acid, oleic acid, linolenic acid and the like; resins obtained by modifying the novolak-type phenolic resin with aromatic hydrocarbons, such as xylene, mesitylene and the like; resins obtained by modifying the novolaktype phenolic resin with rubbers, such as nitrile rubber and the like. These resins are added to the rubber in an amount of 15-45 parts by weight, preferably 20-40 parts by weight, based on 100 parts by weight of carbon 10 black. When the amount of resin is less than 15 parts by weight, the effect of the resin does not substantially appear, while when the amount of resin exceeds 45 parts by weight, excess resin forms agglomerates to cause phase separation in the resulting rubber composition, and deteriorates noticeably the physical properties of the rubber composition.

Further, in the present invention, a mixture of the novolak-type phenolic resin and the novolak-type modified phenolic resin is preferably used, because the use of 20 the mixture can improve synergistically the durability of the resulting rubber composition as compared with the case where these resins are used alone. In this case, the mixing ratio of novolak-type phenolic resin to novolak-type modified phenolic resin, particularly the mixing ratio of novolak-type phenol resin to novolak-type cashew modified phenol resin or to novolak-type tall oil modified phenol resin, shoul be 80/20-20/80, preferably 60/40-40/60.

In the present invention, as the hardener for the resin, aldehyde-donors, that is, aldehyde-generating agents, such as hexamethylenetetramine, paraformaldehyde, hexamethoxymethylmelamine and the like, are preferably used. The hardener is used in an amount enough to 35 harden the resin.

In the present invention, in addition to the above described ingredients, vulcanizing agents, such as sulfur, N,N'-dithiodiamines, thiurums and the like, vulcanization accelerator, antioxidant, fillers other than carbon 40 black, such as silica and the like, process oil and other additives may be contained in the rubber composition.

The following examples are given for the purpose of illustration of this invention and are not intended as limitations thereof.

# EXAMPLE 1

. Homogeneous rubber compositions having a compounding recipe (parts by weight) shown in the following Table 1 were vulcanized at 145° C. for 40 minutes in 50 a press to produce vulcanized rubber sheets having a thickness of 2 mm. The elongation at break (Eb), 20% modulus and dynamic modulus of the rubber sheets are shown in Table 1. The elongation at break and 20% modulus were measured with respect to an ASTM F 55 type dumbbell according to ASTM D 412. The dynamic modulus was measured at room temperature with respect to a strip-shaped sample having a length of 25 mm, a width of 5 mm and a thickness of 2 mm by means of a high-power spectrometer made by Iwamoto 60 Seisakusho by vibrating the sample at a frequency of 10 Hz and under a dynamic strain of 2%, the sample being used under an elongated state of 5% obtained by applying a static pressure.

It can be seen from Table 1 that a rubber composition 65 containing resin and carbon black in a mixing ratio defined in the present invention has remarkably improved 20% modulus and dynamic modulus, and fur-

ther has a satisfactorily high elongation at break for practical use.

In order to make a tire light in weight and to improve the ride feeling thereof, provision was made of a tire 1 shown in FIG. 1, which had a size of 165 SR 13 and comprised a belt layer 2 composed of two steel cord plies and a carcass layer 3 composed of one ply formed of polyethylene terephthalate fiber of 1500 d/2 and a bead filler 6, the carcass ply having a turn-up portion 3' extending up to a low position near a rim flange R<sub>T</sub>. In FIG. 1, the numeral 4 represents a bead portion, the numeral 5 represents a side wall portion, the numeral 7 represents a bead wire, and the letter R represents a rim. Rubber composition No. 1, No. 3, No. 5, No. 7 and No. 8 shown in Table 1 were used as a rubber for the bead filler 6 of the tire shown in FIG. 1, and the high speed performance, lateral rigidity index, ride feeling and durability (condition A) of the tire were evaluated. The obtained results are shown in the following Table 2.

For comparison, rubber composition No. 1 (conventional rubber composition) described in Table 1 was used as a rubber for the bead filler 26 of a conventional tire 21 shown in FIG. 2, which had a structure that the turn-up portion 23' of a carcass ply extended up to the vicinity of the maximum width portion of the side wall portion 25. In FIG. 2, the numeral 22 represents a belt layer, the numeral 23 represents a carcass layer, the numeral 24 represents a bead portion, the numeral 27 represents a bead wire, the numeral 28 represents a cord layer, the letter R represents a rim and the letter RT represents a rim flange. The high speed performance, lateral rigidity index, ride feeling and durability (condition A) of the tire 21 using Rubber composition No. 1 were evaluated. The obtained results are also shown in Table 2.

The evaluation of the performances was carried out in the following manner.

(1) High speed performance:

A tire is assembled in a 4½ J rim, inflated under an internal pressure of 2.1 kg/cm<sup>2</sup> and pressed on a drum having a diameter of 1.7 m under a load of 390 kg/cm<sup>2</sup>. The tire is run on the drum at a speed of 80 km/hr for 2 hours and left to stand for 3 hours. Then, the tire is run at a speed of 121 km/hr for 30 minutes. When the tire runs without trouble, the speed is raised stepwise by 8 km/hr every 30 minutes. The high speed performance of the tire is evaluated by the speed at the breakage of the tire and by the running time until the breakage at the speed.

(2) Lateral rigidity index:

A tire is assembled in a 4½ J rim and inflated under an internal pressure of 1.7 kg/cm<sup>2</sup>. The tire is pressed and fixed to a push car, which has a jagged non-slip surface, under a vertical load of 320 kg, the push car is pulled in a direction perpendicular to the direction of the tire, and the lateral road, which is caused at a lateral shift of the tire of 15 mm, is measured. The lateral rigidity index of a sample tire is indicated by the ratio of the lateral road of the sample tire to that, calculated as 100, of Conventional tire A.

The ride feelings of the above obtained tires were compared with each other by the impact index and damping index measured in the following manner.

'A tire is travelled at a speed of 50 km/hr on a road having rubber projections of 10 mm height fixed thereto, and the vibration subjected to the tire in the up-and-down direction is measured in the form of a

reaction in the rotating shaft of the tire by means of an acceleration meter. The impact absorbing property of a

6 tire without trouble is over 30,000 km, the tire is evaluated as an acceptable tire.

			TA	BLE 1						
Rubber composition No.	1	2	3	4	5	6	7	8	9	10
Compounding recipe (parts by weight)										•
Natural rubber Styrene-butadiene copolymer rubber	100	100	100	100	100	100	100	100	50 50	50
Polybutadiene rubber Carbon black*	70		70	70	70	70	70	100	70	50 70
Novalak-type cashew modified phenol resin**		20	8	14	20	26	40	30	20	20
Stearic acid	2	2	2	2	2	2	2	2	2	2
Zinc white	7.5	7.5	7.5	7.5	7.5	7.5	7.5	7.5	7.5	7.5
N—oxydiethylenebenzothiazole sulfeneamide	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5
Sulfur .	4	4	4	4	4	4	4	4	4	4
Hexamethylenetetramine Property		2	O.8	1.4	2.0	2.6	4.0	3.0	2.0	2.0
Elongation at break (%)	170	380	220	230	225	200	100	145	220	190
20% modulus (kg/cm²)	13.5	7.5	35.0	43.0	60,5	69.0	92.0	89.0	55.0	63.5
Dynamic modulus (kg/cm <sup>2</sup> )	120	40	350	520	730	890	1,530	1,420	710	775

<sup>\*1</sup>A:82 mg/g. DBP:102 ml/100 g \*\*Novoluk-type phenol resin obtained by modifying 100 parts by weight of phenol with 40 parts by weight of cashoew nut oil

	•	TAE	BLE 2			
Tire	Α	В	С	D	E	F
Tire structure Rubber composition used in bead filler Performance	FIG. 2 No. 1	FIG. 1 No. 1	F1G, 1 No. 3	FIG. 1 No. 5	FIG. 1 No. 7	FIG. 1 No. 8
High-speed performance Speed (km/h)		160	4.55	***		-
Running time (min) Lateral rigidity index	185 23 100	169 28	177 27	185 28	193 . 2	193 4
Ride feeling		75	93	103	112	110
Impact index Damping index	100 100	1'15 86	112 93	109 110	105 115	106 113
Durability (condition A)	run over 30,000 km	11,000 km	21,500 km	run over 30,000 km	16,500 km	run over 30,000 km

sample tire is indicated by the impact index, which is the reciprocal of the ratio of the amplitude in the first per- 40 composition of the present invention is used as a bead iod of the above measured wave shape in the tire to that, calculaed as 100, in Conventional tire A.

The vibration damping property of a sample tire is indicated by the damping index, which is the recprocal of the ratio of the damping coefficient calculated from 45 the above measured wave shape in the tire to that, calculated as 100, in Conventional tire A.

# (4) Durability (condition A):

A tire is assembled in a  $4\frac{1}{2}$  J rim, and pressed on a metal drum having a diameter of 1.7 m under an over- 50 load and over internal pressure condition that the strain energy concentrated to the turn-up end of the carcass ply is as large as about 4 times of the strain energy in a practically running tire. Then, the tire is run at a speed of 60 km/hr, and the durability of the tire is shown by 55 the running distance until breakage occurs at the turnup end of carcass ply. When the running distance of a

It can be seen from Table 2 that, when the rubber filler rubber of a tire, the tire has equal or superior to Conventional tire A in the high speed performance, cornering stability and durability and further is remarkably superior to Conventional tire A in the ride feeling.

## EXAMPLE 2

Rubber compositions were produced in the same compounding recipe as that of Rubber composition No. 5 in Table 1, except that only the carbon black is replaced by carbon blacks shown in the following Table 3. The viscosity of the resulting rubber compositions was measured according to JIS K 6300, and the elongation at break, 20% modulus and dynamic modulus thereof were measured in the same manner as described in Example 1. Then, tires having a structure shown in FIG. 1 were produced by the use of the rubber compositions, and the performances of the tires were evaluated in the same manner as described in Example 1. The obtained results are shown in Table 3.

TABLE 3

Rubber Composition No.	· 11	12	13	5	15	16	17			
Carbon black	:									
1A (mg/g)	36	43	86	82 '	84	121	145			
DBP (ml/100 g)	91	· 121	60	102	150	114	113			
Property		12 5/5								
Mooney viscosity	66	73	. 85	92	125	105	132			
Elongation at break (%)	200	220	250	225	200	210	190			
20% modulus (kg/cm <sup>2</sup> )	38.5	50.5	51.5	60.5	45.0	53.0	42.5			

run over 30,000 km

		TABLI	E 3-continue	d	1, 1		
Dynamic modulus (kg/cm²)	420	610	625	750	595	715	530
Tire	G	н	I' ·	D		J	
Performance High-speed performance		1	7 T				
Speed (km/hr) running time (min) Lateral rigidity index	185 10 95	185 21 97	185 20	185 28		185 20	

run over

30,000 km

run over

30,000 km

It can be seen from Table 3 that tires having more improved durability can be obtained by the use of carbon black having an IA of 40-130 mg/g and a DBP of not higher than 130 ml/100 g. Rubber composition No. 15 15 and No. 17 are very poor in the fluidity in the unvulcanized state, and are very difficult in the extrusionmolding. Therefore, the evaluation of tires using the rubber compositions are omitted.

18,000 km

Durability (condition A)

## EXAMPLE: 3

Rubber compositions were produced according to the compounding recipe shown in the following Table 4. The fatigue life of the rubber compositions was measured in the method as explained later, and other prop- 25 shown in Table 5. erties thereof were measured in the same manner as described in Example 1. The obtained results are shown in Table 4.

The fatigue life of the rubber composition was mea-

fatigue tester, and the number of vibrations until the durmbbell was broken was measured.

run over 30,000 km

Then, tires having a structure shown in FIG. 1 were produced by the use of Rubber composition Nos. 22-26 shown in Table 4, and the performance of the tires was evaluated in the same manner as described in Example 1. In this case, the durability of the tires was evaluated, not only under condition A, but also under a more severe condition (condition B), that is, under a super overload and super over internal pressure condition, wherein strain energy concentrated to the turn-up end of carcass ply is as large as 8 times of the strain energy in a practically running tire. The obtained results are

It can be seen from Tables 4 and 5 that, when unmodified phenolic resin is used together with modified phenolic resin, the resulting rubber composition has a synergistically improved durability.

	IABLE 4									
Rubber Composition No.	18	19	20	21	22	23	24	25	26	
Compounding recipe			·							
(parts by weight)							•			
Natural rubber	100	100	100	100	100	100	100	100		
Carbon black*	75	75	75	75	75	75	100 75	100	100	
Cashew-modified phenol resin**	24 -	18	· 18	7.5	13	./3	/3	75	75	
Phenol resin***		. 6	12	. 18	24	••		_		
Tall oil-modified phenol resin****	•		12	. 10 .	24	. 18	12	6		
Stearic acid	•	•	•	_	•	6	12	18	24	
Zine white	10	2	2	2	. 2	2	2	2	· 2	
N—oxydiethylenebenzothiazole	10	10	10	. 10	10	10	10 ·	10	10	
sulfeneamide .	1	· I	1	1 :	1	1	1	1 .	1	
Sulfur		:: :					-		٠.	
	6	6	6	6	6	6 .	6	6	6	
Hexamethylenetetramine	2.4	2.4	2.4	2.4	2.4	2.4	2.4	2,4	2.4	
Property	• •		•	•		. ,			5	
Elongation at break (%).	160	160 ·	155	140	120	· 145	170	165	160	
20% modulus (kg/cm²)	72	70	47 .	50	120	143	170	103	100	

Dynamic modulus (kg/cm²):

\*IA:82 mg/g. DBP:102 ml/100 g

\*Novolak-type phenol resin obtained by modifying 100 parts by weight of phenol with 40 parts by weight of cashew out oil

Fatigue life (number of vibrations)

100 parts by weight of phenol with 40 parts by weight of tall oil

T	nT	*	
ΙΔ	к		

Tire	ĸ	L	М	N	0					
Rubber composition used in bead filler High speed performance	No. 22	No. 23	No. 24	No. 25	No. 26					
Speed (km/h) Running time (min) Lateral rigidity index Durability (condition A)	185 12 98 run over 30,000 km	185 21 101 run over 30,000 km	185 28 105 run over 30,000 km	185 27 106 run over 30,000 km	185 · 28 106 run over 30,000 km					
Durability (condition B)	rubber is broken after 9,800 km running	run over ·· 13,000 km ·	run over 13,000 km :	run over 13,000 km	rubber is broken after 13,000 km running					

sured in the following manner according to ASTM D 65 412. That is, an ASTM F type dumbbell of the rubber composition was fixed under an elongated state of 35% and vibrated at 170° C. by means of an elongation type

What is claimed is:

1. A bead filler rubber composition, consisting essentially of:

(a) 100 parts by weight of a rubber selected from the group consisting of natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene 5 copolymer rubber and blends thereof;

(b) 40-130 parts by weight of carbon black having an iodine adsorbability of 40-130 mg/g and a dibutyl phthalate absorbability of not higher than 130

ml/100 g;

- (c) 15-45 parts by weight per 100 parts by weight of carbon black of a mixture of a novolak phenolic resin selected from the group consisting of novolak phenol resin, novolak cresol resin, novolak resorcinol resin; and a novolak modified phenolic resin obtained by modifying with a compound selected from the group consisting of oils, aromatic hydrocarbons, or rubbers, wherein the mixing ratio of the novolak phenolic resin to the novolak modified phenolic resin is from 80/20 to 20/80 by weight, an 20 effective amount of a hardener for the resin.
- 2. A bead filler rubber composition according to claim 1, wherein the amount of the carbon black is 60-120 parts by weight based on 100 parts by weight of the rubber.
- 3. A bead filler rubber composition according to claim 1, wherein the amount of the novolak-type pheno-

lic resin is 20-40 parts by weight based on 100 parts by weight of the carbon black.

- 4. A bead filler rubber composition according to claim 1, wherein the mixing ratio of the novolak phenolic resin/the modified novolak phenolic resin is 60/40-40/60.
- 5. A bead filler rubber composition according to claim 1 wherein said novolak modified phenolic resins are novolak phenolic resins modified with an oil selected from the group consisting of rosin oil, tall oil and cashew nut oil.
- 6. A bead filler rubber composition according to claim 1, wherein said novolak modified phenolic resins are novolak phenolic resins modified with an oil selected from the group consisting of linoleic acid, oleic acid and linolenic acid.
- 7. A bead filler rubber composition according to claim 1, wherein said novolak modified phenolic resins are novolak phenolic resins modified with an aromatic hydrocarbon selected from the group consisting of xylene and mesitylene.
- 8. A bead filler rubber composition according to claim 1, wherein said novolak modified phenolic resins
  25 are novolak phenolic resins modified with a nitrile rubber.

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# Miyake et al.

Dec. 20, 1983 [45]

[54]	BEAD FIL	LER RUBBER COMPOSITION
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[73]	Assignee:	Bridgestone Tire Company Limited, Tokyo, Japan
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Oct	. 18, 1978 <b>[J</b> f	P] Japan 53-127256
[51] [52]	U.S. Cl	
[58]	Field of Sea	arch 524/495, 511; 525/139; 152/362 R; 260/727, 775, 779 R

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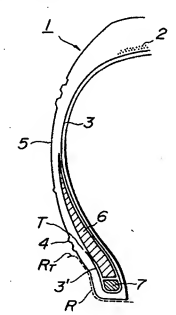
Primary Examiner-Theodore E. Pertilla Attorney, Agent, or Firm-Sughrue, Mion, Zinn, Macpeak, and Seas

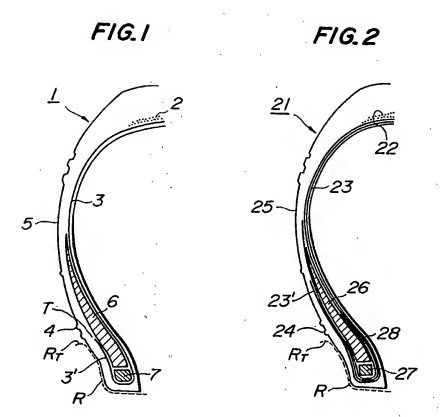
# [57]

## **ABSTRACT**

A rubber composition comprising rubber, novolak-type phenolic resin and carbon black is very rigid and is adapted to be used as a bead filler rubber of a tire. Tires using the rubber composition are excellent in the high speed performance, lateral rigidity, ride feeling and durability.

# 8 Claims, 2 Drawing Figures





# BEAD FILLER RUBBER COMPOSITION

This is a continuation of application Ser. No. 85,346, filed Oct. 16, 1979, now abandoned.

# BACKGROUND OF THE INVENTION

#### 1. Field of the Invention

The present invention relates to a rubber composition, and more particularly, relates to a super rigid rub- 10 ber composition comprising rubber, novolak-type phenolic resin and carbon black, and adapted to be used as a bead filler rubber of a tire.

# 2. Description of the Prior Art

There have been variously investigated the structure 15 have accomplished the present invention. of bead portion of radial tire in order to satisfy the rigidity and durability required to tire. For example, it has been attempted to improve the dynamic performance and durability of tire by arranging a bead-reinforcing layer in the bead portion. However, this method 20 has drawbacks that a large number of production steps are necessary and the productivity of the tire is very

Japanese Utility Model Application No. 16,084/72, French Patent No. 1,260,138 and U.S. Pat. No. 25 4,067,373 disclose methods for improving the running performance and other property by arranging super rigid rubber in the bead portion. However, these methods do not substantially think of rubber, which can develop fully a function as a bead filler rubber subjected 30 to complicated forces during the running of a tire and can give a sufficiently high durability to a rubber tire.

While, it is well known to use novolak-type phenolic resin in the production of rigid rubber compositions. However, these rubber compositions substantially con- 35 cern nitrile rubber and neoprene rubber having a high compatibility with the resin. The nitrile rubber series of neoprene rubber series rigid rubber is very difficult to be vulcanized together with natural rubber, polybutadiene rubber and the like, which are commonly used as a 40 rubber for tire, and therefore when the nitrile rubber series or neoprene rubber series rigid rubber is used as a tire part, the rubber is apt to be separated from natural rubber, polybutadiene rubber or the like, and can not be practically used.

In order to solve the above described drawbacks, the inventors have variously investigated how to produce a super rigid rubber composition by compounding novolak-type phenolic resin to natural rubber, polybutadiene rubber or the like, and found out the following facts. 50 Novolak-type phenolic resin is essentially incompatible with natural rubber and other rubbers. Therefore, novolak-type phenolic resin is formed into spherical agglomerates during the kneading commonly carried out in the production of rubber, and has the filling effect only. 55 This phenomenon still occurs even when various resins are used as a resin in place of novolak-type phenolic resin for producing a homogenous mixture of the resin with the rubber. However, when novolak-type phenolic resin is compounded to natural rubber or other rubber 60 together with carbon black, the mixture of the resin and carbon black exhibits a reinforcing effect on the rubber, which is completely different from the reinforcing effect of the resin alone on the rubber or the reinforcing effect of the carbon black alone on the rubber, depend- 65 ing upon the mixing ratio of the resin to the carbon black, and a super rigid rubber composition having a durability remarkably superior to that of conventional

resin-reinforced rubber can be obtained. That is, when natural rubber or diene series rubber is mixed with novolak-type phenolic resin, which is essentially incompatible with the rubber, the resin is separated from the rubber to form island-like large agglomerates in the mixture, but when a mixture of novolak-type phenolic resin and carbon black is compounded to natural rubber or other diene series rubber, the resin disperses uniformly in the rubber without forming large agglomerates similarly to the dispersed state in an ordinary mixture of rubber and carbon black. This action of carbon black to the resin is highly influenced by the mixing ratio of the carbon black to the resin and by the kind of the carbon black. Based on the discovery, the inventors

#### SUMMARY OF THE INVENTION

The feature of the present invention is the provision of a bead filler rubber composition, comprising 100 parts by weight of a rubber selected from the group consisting of natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene copolymer rubber and blends thereof, 40-130 parts by weight of carbon black, 15-45 parts by weight based on 100 parts by weight of the carbon black of at least one of novolaktype phenolic resin and novolak-type modified phenolic resin, and a hardener for the resin.

# BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a cross-sectional view of the side portion of a tire according to the present invention; and FIG. 2 is a cross-sectional view of the side portion of a conventional tire.

## DESCRIPTION OF THE PREFERRED **EMBODIMENTS**

The rubber to be used in the present invention includes natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene copolymer rubber and blends thereof. Carbon black is compounded to the rubber in an amount of 40-130 parts by weight, preferably 60-120 parts by weight, more preferably 65-85 parts by weight, based on 100 parts by weight of the rubber. The use of carbon black in an amount of less than 40 parts by weight is too small to disperse homogeneously the necessary amount of resin for reinforcing the rubber. While, when the amount of carbon black is more than 130 parts by weight, the resulting rubber composition is brittle and is very poor in the durability. Further, in the present invention, carbon black having an iodine adsorbability (IA) of 40-130 mg/g and a dibutyl phthalate absorbability (DBP) of not more than 130 ml/100 g defined in ASTM D 1765 is preferably used. Carbon black having an IA of less than 40 mg/g affects adversely the dispersibility of the resin, and carbon black having an IA of more than 130 mg/g is poor in the dispersibility in itself. Therefore, the use of such carbon black is not preferable. When carbon black has a DBP of more then 130 ml/100 g, the carbon black used in an amount sufficient to disperse the resin can not be fully dispersed in the rubber.

In the present invention, the above described carbon black is used together with at least one of novolak-type phenolic resin and novolak-type modified phenolic resin. The novolak-type phenolic resin includes novolak-type phenol resin, novolak-type cresol resin and novolak-type resorcinol resin. The novolak-type modified phenolic resin includes resins obtained by modify-

ing the above described novolak-type phenolic resin with oils, such as rosin oil, tall oil, cashew nut oil, linoleic acid, oleic acid, linolenic acid and the like; resins obtained by modifying the novolak-type phenolic resin with aromatic hydrocarbons, such as xylene, mesitylene 5 and the like; resins obtained by modifying the novolaktype phenolic resin with rubbers, such as nitrile rubber and the like. These resins are added to the rubber in an amount of 15-45 parts by weight, preferably 20-40 parts by weight, based on 100 parts by weight of carbon 10 black. When the amount of resin is less than 15 parts by weight, the effect of the resin does not substantially appear, while when the amount of resin exceeds 45 parts by weight, excess resin forms agglomerates to cause phase separation in the resulting rubber composition, 15 and deteriorates noticeably the physical properties of the rubber composition.

Further, in the present invention, a mixture of the novolak-type phenolic resin and the novolak-type modified phenolic resin is preferably used, because the use of 20 the mixture can improve synergistically the durability of the resulting rubber composition as compared with the case where these resins are used alone. In this case, the mixing ratio of novolak-type phenolic resin to novolak-type modified phenolic resin, particularly the mixing ratio of novolak-type phenol resin to novolak-type cashew modified phenol resin or to novolak-type tall oil modified phenol resin, shoud be 80/20-20/80, preferably 60/40-40/60.

In the present invention, as the hardener for the resin, aldehyde-donors, that is, aldehyde-generating agents, such as hexamethylenetetramine, paraformaldehyde, hexamethoxymethylmelamine and the like, are preferably used. The hardener is used in an amount enough to 35 harden the resin.

In the present invention, in addition to the above described ingredients, vulcanizing agents, such as sulfur, N,N'-dithiodiamines, thiurums and the like, vulcanization accelerator, antioxidant, fillers other than carbon 40 black, such as silica and the like, process oil and other additives may be contained in the rubber composition.

The following examples are given for the purpose of illustration of this invention and are not intended as limitations thereof.

# EXAMPLE 1

Homogeneous rubber compositions having a compounding recipe (parts by weight) shown in the following Table 1 were vulcanized at 145° C. for 40 minutes in 50 a press to produce vulcanized rubber sheets having a thickness of 2 mm. The elongation at break (Eb), 20% modulus and dynamic modulus of the rubber sheets are shown in Table 1. The elongation at break and 20% modulus were measured with respect to an ASTM F 55 type dumbbell according to ASTM D 412. The dynamic modulus was measured at room temperature with respect to a strip-shaped sample having a length of 25 mm, a width of 5 mm and a thickness of 2 mm by means of a high-power spectrometer made by Iwamoto 60 Seisakusho by vibrating the sample at a frequency of 10 Hz and under a dynamic strain of 2%, the sample being used under an elongated state of 5% obtained by applying a static pressure.

It can be seen from Table 1 that a rubber composition 65 containing resin and carbon black in a mixing ratio defined in the present invention has remarkably improved 20% modulus and dynamic modulus, and fur-

ther has a satisfactorily high elongation at break for practical use.

In order to make a tire light in weight and to improve the ride feeling thereof, provision was made of a tire 1 shown in FIG. 1, which had a size of 165 SR 13 and comprised a belt layer 2 composed of two steel cord plies and a carcass layer 3 composed of one ply formed of polyethylene terephthalate fiber of 1500 d/2 and a bead filler 6, the carcass ply having a turn-up portion 3' extending up to a low position near a rim flange RT. In FIG. 1, the numeral 4 represents a bead portion, the numeral 5 represents a side wall portion, the numeral 7 represents a bead wire, and the letter R represents a rim. Rubber composition No. 1, No. 3, No. 5, No. 7 and No. 8 shown in Table 1 were used as a rubber for the bead filler 6 of the tire shown in FIG. 1, and the high speed performance, lateral rigidity index, ride feeling and durability (condition A) of the tire were evaluated. The obtained results are shown in the following Table 2.

For comparison, rubber composition No. 1 (conventional rubber composition) described in Table 1 was used as a rubber for the bead filler 26 of a conventional tire 21 shown in FIG. 2, which had a structure that the turn-up portion 23' of a carcass ply extended up to the vicinity of the maximum width portion of the side wall portion 25. In FIG. 2, the numeral 22 represents a belt layer, the numeral 23 represents a carcass layer, the numeral 24 represents a bead portion, the numeral 27 represents a bead wire, the numeral 28 represents a cord layer, the letter R represents a rim and the letter RT represents a rim flange. The high speed performance, lateral rigidity index, ride feeling and durability (condition A) of the tire 21 using Rubber composition No. 1 were evaluated. The obtained results are also shown in

The evaluation of the performances was carried out in the following manner.

(1) High speed performance:

A tire is assembled in a 4½ J rim, inflated under an internal pressure of 2.1 kg/cm<sup>2</sup> and pressed on a drum having a diameter of 1.7 m under a load of 390 kg/cm<sup>2</sup>. The tire is run on the drum at a speed of 80 km/hr for 2 hours and left to stand for 3 hours. Then, the tire is run at a speed of 121 km/hr for 30 minutes. When the tire 45 runs without trouble, the speed is raised stepwise by 8 km/hr every 30 minutes. The high speed performance of the tire is evaluated by the speed at the breakage of the tire and by the running time until the breakage at the speed.

(2) Lateral rigidity index:

A tire is assembled in a 4½ J rim and inflated under an internal pressure of 1.7 kg/cm<sup>2</sup>. The tire is pressed and fixed to a push car, which has a jagged non-slip surface, under a vertical load of 320 kg, the push car is pulled in a direction perpendicular to the direction of the tire, and the lateral road, which is caused at a lateral shift of the tire of 15 mm, is measured. The lateral rigidity index of a sample tire is indicated by the ratio of the lateral road of the sample tire to that, calculated as 100, of Conventional tire A.

(3) Ride feeling:

The ride feelings of the above obtained tires were compared with each other by the impact index and damping index measured in the following manner.

A tire is travelled at a speed of 50 km/hr on a road having rubber projections of 10 mm height fixed thereto, and the vibration subjected to the tire in the up-and-down direction is measured in the form of a

reaction in the rotating shaft of the tire by means of an acceleration meter. The impact absorbing property of a

6 tire without trouble is over 30,000 km, the tire is evaluated as an acceptable tire.

		TABLE 1									
Rubber composition No.	1	2	3	4	5	6	7	8	9	10	
Compounding recipe (parts by weight)		-						·	· · · · · · · · · · · · · · · · · · ·	-	
Natural rubber Styrene-butadiene copolymer rubber Polybutadiene rubber	100	100	100	100	100	100	100	100	50 50	50 50	
Carbon black* Novalak-type cashew modified phenol resin**	70	20	<b>7</b> 0 8	70 14	70 20	70 26	70 40	100 30	70 20	70 20	
Stearic acid Zinc white	2 7.5	2	2	2	2	2	2	2	2	2	
N—oxydiethylenebenzothiazole sulfeneamide	1.5	7.5 1.5	7.5 1.5	7.5 1.5	7.5 1.5	7.5 1.5	7.5 1.5	7.5 · 1.5	7.5 1.5	7.5 1.5	
Sulfur .	4	4	4	4	4	4	4	4	4	. 4	
Hexamethylenetetramine Property		2	0.8	1.4	2.0	2.6	4.0	3.0	2.0	2.0	
Elongation at break (%) 20% modulus (kg/cm²) Dynamic modulus (kg/cm²)	170 13.5 120	380 7.5 40	220 35.0 350	230 43.0 520	225 60.5 730	200 69.0 890	100 92.0 1,530	145 89.0 1,420	220 55.0 710	190 63.5 775	

<sup>\*1</sup>A:82 mg/g. DBP:102 ml/100 g

Novoluk-type phenol resin obtained by modifying 100 parts by weight of phenol with 40 parts by weight of cashoew nut oil

		TAI	BLE 2		•	
Tire	A	В	c ·	Ď	E	F
Tire structure Rubber composition used in bead filler Performance High-speed performance	FIG. 2 No. 1	FIG. 1 No. 1	FIG. 1 No. 3	FIG. 1 No. 5	FIG. 1 No. 7	FIG. 1 No. 8
Speed (km/h) Running time (min) Lateral rigidity index Ride feeling	185 23 100	169 28 75	177 27 93	185 . 28 103	193 - 2 112	193 4 110
Impact index Damping index Durability (condition A)	100 100 run over 30,000 km	115 86 11,000 km	112 93 21,500 km	109 110 run over 30,000 km	105 115 16,500 km	106 113 run over 30,000 km

sample tire is indicated by the impact index, which is the reciprocal of the ratio of the amplitude in the first per- 40 composition of the present invention is used as a bead iod of the above measured wave shape in the tire to that, calculaed as 100, in Conventional tire A.

The vibration damping property of a sample tire is indicated by the damping index, which is the recprocal of the ratio of the damping coefficient calculated from 45 the above measured wave shape in the tire to that, calculated as 100, in Conventional tire A.

# (4) Durability (condition A):

A tire is assembled in a 4½ J rim, and pressed on a metal drum having a diameter of 1.7 m under an over- 50 load and over internal pressure condition that the strain energy concentrated to the turn-up end of the carcass ply is as large as about 4 times of the strain energy in a practically running tire. Then, the tire is run at a speed of 60 km/hr, and the durability of the tire is shown by 55 the running distance until breakage occurs at the turnup end of carcass ply. When the running distance of a

It can be seen from Table 2 that, when the rubber filler rubber of a tire, the tire has equal or superior to Conventional tire A in the high speed performance, cornering stability and durability and further is remarkably superior to Conventional tire A in the ride feeling.

## EXAMPLE.2

Rubber compositions were produced in the same compounding recipe as that of Rubber composition No. 5 in Table 1, except that only the carbon black is replaced by carbon blacks shown in the following Table 3. The viscosity of the resulting rubber compositions was measured according to JIS K 6300, and the elongation at break, 20% modulus and dynamic modulus thereof were measured in the same manner as described in Example 1. Then, tires having a structure shown in FIG. 1 were produced by the use of the rubber compositions, and the performances of the tires were evaluated in the same manner as described in Example 1. The obtained results are shown in Table 3.

~ 4	TOT	$\overline{}$	•
IΑ	BI		•

		IAL	SLE 3	•			
Rubber Composition No.	11	12	13	5	15	16	17
Carbon black	•						
1A (mg/g)	36	43	86	82 .	84	121	145
DBP (ml/100 g)	91	· 121 ·	. 60	102	150	114	113
Property							
Mooney viscosity	66	73	85	92	125	105	132
Elongation at break (%)	200	220	250	225	200	210	190
20% modulus (kg/cm <sup>2</sup> )	38.5	50.5	51.5	60.5	45.0	53.0	42.5

ТΔ	RI	F	2	tinued	

			2 Continue				
Dynamic modulus (kg/cm²)	420	610	625	750	595	715	530
Tire	G	Н	ľ			J	
Performance High-speed performance			7 Y		· ·		
Speed (km/hr) running time (min) Lateral rigidity index Durability (condition A)	185 10 95 18,000 km	185 21 97 run over	185 20 97 run over	185 28 103 run over		185 20 98 run over	
		30,000 km	30,000 km	30,000 km		30,000 km	

It can be seen from Table 3 that tires having more improved durability can be obtained by the use of carbon black having an IA of 40-130 mg/g and a DBP of not higher than 130 ml/100 g. Rubber composition No. 15 15 and No. 17 are very poor in the fluidity in the unvulcanized state, and are very difficult in the extrusionmolding. Therefore, the evaluation of tires using the rubber compositions are omitted.

## EXAMPLE 3

Rubber compositions were produced according to the compounding recipe shown in the following Table 4. The fatigue life of the rubber compositions was measured in the method as explained later, and other prop- 25 erties thereof were measured in the same manner as described in Example 1. The obtained results are shown in Table 4.

The fatigue life of the rubber composition was mea-

fatigue tester, and the number of vibrations until the durmbbell was broken was measured.

Then, tires having a structure shown in FIG. 1 were produced by the use of Rubber composition Nos. 22-26 shown in Table 4, and the performance of the tires was evaluated in the same manner as described in Example 1. In this case, the durability of the tires was evaluated, not only under condition A, but also under a more severe condition (condition B), that is, under a super overload and super over internal pressure condition, wherein strain energy concentrated to the turn-up end of carcass ply is as large as 8 times of the strain energy in a practically running tire. The obtained results are shown in Table 5.

It can be seen from Tables 4 and 5 that, when unmodified phenolic resin is used together with modified phenolic resin, the resulting rubber composition has a synergistically improved durability.

TABLE 4

			*****	<b>-</b> .					
Rubber Composition No.	18	19	20	21	22	23	24	25	26
Compounding recipe (parts by weight)			<del></del> ,						
Natural rubber Carbon black*	100	100	100	100	100	100	100	100	100
Cashew-modified phonol resin**	75 24	75 18	. 18	75 6	75	.75	75	75	75
Phenol resin*** Tall oil-modified phenol resin****		6 .	12	. 18 .	24	18	12	6	
Stearic acid	. 2	. 2	2	2	2	6 2	12	18	24
Zinc white N—oxydiethylenebenzothiazole	10	10	10	10	10	10	10	10	ΙŌ
sulfeneamide Sulfur	•				. •	. 1	i	I	1
Hexamethylenetetramine Property	2.4	6 2.4	6 2,4	6 2.4	6 2.4	. 6, . 2.4	6 2.4	6 2.4	. 6 2.4
Elongation at break (%)	160 72	160	155	140	120	145	170	165	160
Dynamic modulus (kg/cm²)  Fatigue life (number of vibrations)	980 2 × 10 <sup>6</sup>	70 920 3 × 10 <sup>6</sup>	67 : 890 5 × 10 <sup>6</sup>	58 · · · 7t5 . 4 × 106	52 650 3 × 10 <sup>6</sup>	58 710 6 × 10 <sup>6</sup>	65 780 8 × 10 <sup>6</sup>	66 840 4 × 10 <sup>6</sup>	67 880 1 × 10 <sup>6</sup>

•1A:82 mg/g. DBP:102 mt/100 g
•Novolak-type phenol resin obtained by modifying 100 parts by weight of phenol with 40 parts by weight of cashew nut oil
••••Unmodified novolak-type phenol resin
••••Unmodified novolak-type phenol resin

Novolak-type phenol resin obtained by modifying 100 parts by weight of phenol with 40 parts by weight of tall oil

		IABLE	• • •		•
Tire	ĸ	· L	М	N ·	0
Rubber composition used in bead filler High speed performance	No. 22	No. 23	No. 24	No. 25	No. 26
Speed (km/h) Running time (min) Lateral rigidity index	185 12 98	185 21 101	185 28 105	185 27 106	185 · 28 106
Durability (condition A)  Durability (condition B)	run over 30,000 km rubber is broken after 9,800 km running	run over 30,000 km run over ··· 13,000 km	run over 30,000 km run over 13,000 km	run over 30,000 km run over 13,000 km	run over 30,000 km rubber is broken after 13,000 km running

sured in the following manner according to ASTM D 65 412. That is, an ASTM F type dumbbell of the rubber composition was fixed under an elongated state of 35% and vibrated at 170° C. by means of an elongation type

What is claimed is:

- 1 A bead filler rubber composition, consisting essentially of:
  - (a) 100 parts by weight of a rubber selected from the group consisting of natural rubber, polyisoprene rubber, polybutadiene rubber, styrene-butadiene 5 copolymer rubber and blends thereof;

(b) 40-130 parts by weight of carbon black having an iodine adsorbability of 40-130 mg/g and a dibutyl phthalate absorbability of not higher than 130 ml/100 g;

- (c) 15-45 parts by weight per 100 parts by weight of carbon black of a mixture of a novolak phenolic resin selected from the group consisting of novolak phenol resin, novolak cresol resin, novolak resorcinol resin; and a novolak modified phenolic resin 15 obtained by modifying with a compound selected from the group consisting of oils, aromatic hydrocarbons, or rubbers, wherein the mixing ratio of the novolak phenolic resin to the novolak modified phenolic resin is from 80/20 to 20/80 by weight, an 20 effective amount of a hardener for the resin.
- 2. A bead filler rubber composition according to claim 1, wherein the amount of the carbon black is 60-120 parts by weight based on 100 parts by weight of the rubber.
- 3. A bead filler rubber composition according to claim 1, wherein the amount of the novolak-type pheno-

lic resin is 20-40 parts by weight based on 100 parts by weight of the carbon black.

- 4. A bead filler rubber composition according to claim 1, wherein the mixing ratio of the novolak phenolic resin/the modified novolak phenolic resin is 60/40-40/60.
- 5. A bead filler rubber composition according to claim 1 wherein said novolak modified phenolic resins are novolak phenolic resins modified with an oil selected from the group consisting of rosin oil, tall oil and cashew nut oil.
- 6. A bead filler rubber composition according to claim 1, wherein said novolak modified phenolic resins are novolak phenolic resins modified with an oil selected from the group consisting of linoleic acid, oleic acid and linolenic acid.
- carbons, or rubbers, wherein the mixing ratio of the novolak phenolic resin to the novolak modified phenolic resin is from 80/20 to 20/80 by weight, an 20 effective amount of a hardener for the resin.

  A bead filler rubber composition according to 1. A bead filler rubber composition according to 20 elaim 1, wherein said novolak modified phenolic resins are novolak phenolic resins modified with an aromatic hydrocarbon selected from the group consisting of xylene and mesitylene.
  - A bead filler rubber composition according to claim 1, wherein said novolak modified phenolic resins
     are novolak phenolic resins modified with a nitrile rubber.

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# ◎ 特 許 公 報 (B2) 昭57-30856

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## 毎ビードフィラーゴム組成物

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#### 釣特許請求の範囲

2 特許請求の範囲第1項記載のゴム組成物にお ーゴムにいて、カーボンブランクがヨウ素吸着量(IA)が 30 関する。 40~130 刷/3、ジブチルフタレート吸油量 ラジブ (DBP)が130 和/1909以下であるカーボンブ 要求され ランクであるビードフイラーゴム組成物。 検討がた

3 特許請求の範囲第1項記載のゴム組成物にお ド部に配置することによつてタイヤの運動性能、 いて、カーボンブラツクの配合量が、ゴム100 35 耐久性能等を改善させる試みがなされているか、

重量部に対して60~120重量部であるビード フイラーゴム組成物。

4 特許請求の範囲第1項記載のゴム組成物において、ノボラツク型フェノール系樹脂の配合量が5 カーボンブラツク100 賃量部当り20~40重量部であるビードフイラーゴム組成物。

5 特許請求の範囲第1項記載のゴム組成物において、ノボラツク型フエノール系術脂がノボラツク型フエノール樹脂、ノボラツク型クレゾール樹脂、ノボラツク型レゾルシン樹脂であるビードフィラーゴム組成物。

6 特許請求の範囲第1項記載のゴム組成物において、ノボラツク型変性フェノール系樹脂が、ロジン油、トール油、カシュー油、リノール酸、オ15 レイン酸、リノレイン酸等のオイルで変性したノボラツク型フェノール系樹脂、キシレン、メシチレン等の芳香族炭化水素で変性したノボラツク型フェノール系樹脂、ニトリルゴム等のゴムで変性したノボラック型フェノール系樹脂であるビード2007イラーで4個財物

7 特許請求の範囲第1項記載のゴム組成物において、ノボラツク型フェノール系樹脂とノボラツク型変性フェノール系樹脂の混合割合が80/20~20/80であるビードフィラーゴム組成物。

本発明はゴム組成物、詳しくはノボラツク型フェノール系樹脂とカーボンブラツクを併用して配合することによつて、特にタイヤのビードフィラーゴムに好適に使用される整硬質のゴム組成物に関する。

ラジアルタイヤのビード部構造はタイヤとして 要求される関性と耐久性を満足させるべく種々の 検討がなされており、例えばビード補強層をビー ド部に配置することによつてタイヤの運動性能、

このようなものは製造工数が多くなり生産性が著 しく劣るといつた欠点を有していた。

一方、超硬質のゴムをビード部に配置すること によって運動性能等を改善することは、実公昭 47-16084号公報、フランス園粉許第 1,260.138号明細書、米国特許第4,067,373 号明細書等によりよく知られていることである。 しかしタイヤ走行中の複雑な入力下にあるビード フイラーゴムとしての機能を十分に発揮させ、そ んど考慮されていないのが現状である。

他方、硬質ゴム翅成物としては、ノボラツク型 フエノール系樹脂を使用することは以前からよく 知られているが、それは歯脂と相密性の良いニト リルゴムやネオプレンゴムについてのものがほと 15 んどで、このようなニトリルゴムやネオプレンゴ ム系の硬質ゴムをタイヤ部材として使用するには、 従来からタイヤ用ゴムとしてよく使用されている 天然ゴム、ポリブタジェンゴム等との共加硫性が 用に供し得ない。

上記欠点を解決するため本発明者らは天然ゴム やポリプタジエンゴム等にノボラツク型フエノー ル系樹脂を配合して超硬ゴム組成物を得んとして 種々研究したところ、天然ゴム等とノボラツク型 25 の耐久性が極端に悪化するため好ましくない。 更 フェノール系樹脂は本質的に非相密であり、通常 のゴム製造で行なわれているゴム混練が手法では ノボラツク型フェノール系樹脂は球状の凝集体と なつてしまい、単なる光順効果しか示さず、との 樹脂を用いたとしても大差はない。しかしながら 上記の天然ゴム等とノボラツク型フェノール系樹 脂の茶にカーボンブラツクを併用するとその併用 比により、ゴムと樹脂系による補造効果あるいは く異質の補強効果が現われ、従来の樹脂精強ゴム には見られなかつた極めて耐久性に優れた超硬ゴ ム組成物が得られることを確かめた。すなわち本 釆互に非相密で梅島状の大きな相分離状態となる 天然ゴムや他のジエン系ゴムとノボラツク型フエ 40 するが、ノボラツク型フエノール系樹脂とは、ノ ノール系樹脂との混合系にカーボンブラツクを併 用していくと、ゴム中で樹脂は大きな相分離状態 から通常のゴムとカーポンプラツク系にみられる 様なカーボンブラツクの分散状態に近い領域まで

均質に分散されるようになり、このカーボンブラ ツクの樹脂に対する相互作用はカーボンブラツク と側盾の併用比あるいはカーポンプラツクの種類 によつても大きく影響を受けることを見出し本発 5 男を達成するに至つた。

従つて本発明は、天然ゴム、ポリイソプレンゴ ム、ポリプタジエンゴムまたはスチレンープタジ エン共重合体ゴム若しくはとれらのプレンドゴム 100 重量部に対して、40~130重量部のカ の上でタイヤとして必要な耐久性等についてほと 10 ーポンプラツクと、カーボンプラツク100重量 部当り15~45重量部の割合でノボランク型フ エノール系樹脂および/またはノボラツク型変性 フエノール系樹脂を加え、更に樹脂用硬化剤を配 合したビードフイラーゴム組成物に関する。

本発明において使用するゴムは上記の如く天然 ゴム、ポリイソプレンゴム、ポリブタジエンゴム、 またはスチレンープタジエン共重合体ゴム若しく はこれらのブレンドゴムで、かかるゴム100盒 盤部に対してカーボンプラツクを40~130簸 著しく劣るため、セパレーション等を転じ易く奥 20 量部好ましくは60~120重量部、更に好まし くは65~85重量部配合する。カーボンブラツ クが40重量部以下では、積強効果を与えるのに 必要な量の倒脂を均一に分配させるには少なすぎ、 130重量部以上ではゴムが脆くなりゴム組成物 に本発明ではカーボンブラツクとしてASTMD 1765に規定するヨウ素吸着量(IA)が40~ 1 3 0 m/ 8、ジブチルフタレート吸油量(DBP) が130吨/1008以下のカーポンプラツクであ ことはゴムとの相溶性を高めるために各種の変性 30 ればより好適である。IAが40㎏/9以下では 樹脂の分散性に悪影響を与え、130%/1008 以上ではカーボンブラツク自体の分散性が劣るた め好ましくない。またDBPが130me/1009 以上では、樹脂を分散させるために必要なカーボ ゴムとカーボンブラツク系による補強効果とは全 35 ンプラツク量において、カーボンブラツクの分散 性が悪化してしまう。

> 本発明の組成物においては、上記カーボンブラ ツクに、ノボラツク型フェノール系機能および/ またはノボラツク型変性フェノール系制脂を併用 ボラツク型フエノール樹脂、ノボラツク型クレゾ ール歯脂、ノボラツク型レゾルシン齒脂であり、 ノボラツク型変性フエノール系樹脂とは前記のノ ボラツク型フエノール系数艦を、ロジン油、トー

ル油、カシユー油、リノール酸、オレイン酸。**リ** ノレイン酸等のオイルで変性した樹脂、キシレン、 メシチレン等の芳香族炭化水素で変性した樹脂、 ニトリルゴム等のゴムで変性した樹脂である。そ してとれらの樹脂はカーボンブラツク100重量 5 で測定した。 部当り15~45貫量部、好ましくは20~40 重量部をゴムに加える。樹脂の添加量が15重量 部以下では樹脂を添加した効果がほとんどなく補 強効果は期待できず、4.5重量部以上では過剰の 樹脂が凝集体を形成し相分離を起とし、ゴム組成 10 している。 物の物機を響しく低下させるため好ましくない。 また本発明においては後述の実施例で説明するよ うにノボラツク型フェノール系樹脂とノボラツク 型変性フェノール系樹脂を併用することによつて ムの耐久性が相乗的に向上するため、このように 併用して用いることが好ましい。この場合、ノボ ラツク型フエノール系樹脂とノボラツク型変性フ エノール系樹脂、好ましくはノボラツク型フェノ 歯脂あるいはノボラツク型トール油変性フエノー ル樹脂を併用した時の混合割合は重量比で80/ 20-20/80、更に好ましくは60/40~ 40/60である。

供与体であるヘキサメチレンテトラミン、バラホ ルムアルデヒド、ヘキサメトキシメチルメラミン 等のホルムアルデヒド発生剤が好適で、個脂を硬 化させるのに必要な量を配合することができる。

本発明においては、前記以外にも確費、N,N 30 1) 高速走行性 ージチオジアミン類やチウラム類等の加硫剤、加 硫促進剤、老化防止剤、カーボンブラツク以外の 例えばシリカ等の充填剤、プロセスオイル等を添 加しても良い。

以下契施例により本発明を更に詳しく説明する。35 実施例 1

第1聚に示す配合内容(重量部)により、混練 りした各種ゴム組成物をプレス中にて145℃で 4 0 分間加硫して、厚さ 2 mmの加硫ゴムシートを 作成した。このゴムシートを用いて破断時の停び 40 2) 横剛性指数 (Eb)、20%モジュラスおよび動的弾性率を測 定した。結果を解1表に示す。尚、Ebと20% モジュラスについてはASTMD 412に準じ ASTM P型ダンベルにて測定した。また動的弾性

率については岩本製作所製ハイパワースペクトロ メーターを用い、室弧にて長さ25mm、幅5mm、 厚さ2㎞の短冊状サンプルに、静的に5%伸長さ せた状態で、周波数10Hz、動歪2%で振動させ

第1表より明らかな如く、樹脂とカーボンブラ ツクを本発明の併用比にて配合したゴム組成物は 20%モジュラスおよび動的弾性率が著しく向上 し、破断伸びにおいても実用上満足される値を有

次いでタイヤの軽量化や乗心地性能を向上させ るために、顔1図に示すようなベルト騰2として 2プライのスチールコード階とカーカス層3とし て1500 4/2 の ポリエテレンテレフタレート 各々単独で使用した場合に此較して、添加したゴ 15 繊維からなる際の1プライを備えたサイズ1 6 5 SR 13のダイヤ1において、カーカスプライの 折返し 3をリムフランジ fl r の近傍の低い位置に とどめた供試タイヤ1のビードフィラー6のゴム として前記第1表に示す配合ル1,ル3,ル5, ール樹脂とノボラツク型カシユー変性フエノール 20 私7および私8の各種ゴム組成物を用いて高速定 行性、機剛性指数、特殊耐久ドラム試験(A条件、 B条件)について検討した。結果を第2表に示す。 参考のために、第2回に示すカーカスプライの折 返し23をサイドウオール部25の最大幅付近ま 本発明において、樹脂用硬化剤とはアルデヒド 25 で延長した構造を省するタイヤ21のビードフィ ラー26のゴムとして、前記第1表の配合※1の ゴム組成物を用いた従来タイヤについても検討し、 結果を合せて第2表に示す。

尚、評価法は次の通りである。

タイヤを4 🛂 Jリムに組み、 2.1 kg/ciiの 内圧を充塡し、直径 1.7mの金属ドラム上に 390kgの荷重で圧然し、80㎞/hの速度で 2時間の俄し走行を行なった後、3時間放置。 次いで121㎞/hの速度で30分間走行させ 異常なく完定すると、8km/h速度を上げ30 分間走行させる。同様に完走すると以下8㎞/h 30分の刻みで連続ステツプアップし、放瞳発 生時の速度および走行時間を評価した。

タイヤを4 シッよりムに組み1.7kg/cifの内 圧を充填し、表面に齷歯状の滑り止めを施した 台車上に320kgの垂直荷重で圧着固定し、タ イヤの側方向に台車を引張り、タイヤ横変位

15mの時に生じる横荷重を測定し従来タイヤを100として指数で表示した。

## 3) 兼心地性

路面に高さ10mmのゴム状突起を固定し、こ イのの上を50mm/hの速度で通過するときの上下 5 通常方向振動をタイヤ取付軸の反力として加速度計 の条にて測定した。この時記録される波形から第1 60 関期の振幅を当り指数として、従来タイヤ対比 が返めで表示、又、上記の記録液形より減衰係 示し数を求め、従来タイヤ対比の遊動の減衰指数と 10 た。

8

して餐示した。

# 4) 特殊耐久ドラム試験(A条件)

タイヤを4 1/2 J リムに組み、カーカスプライの折返し端部に集中する歪エネルギーが実車 通常走行時の約 4倍となる様な過荷度、過内圧の条件で直径 1.7 mの金属 ドラム上に圧着して60㎞/nの速度で走行させ、カーカスプライ折返し端部に故障が発生するまでの走行距離で示したが30,000㎞まで走行すれば合格とした。

榌

뫲

٥	•	<b>\$ 6</b>	ء ۾	4 F.	<del>رب</del> ري	ري د د د	₹ 1
1	40	<i>∞</i>	63	<b></b>			19
σ	9 9	© (~	20	4 G÷	1,5	4 %	2 2 6 5 5, 0 7 1 0
బ	100	160	3.0	7 <del>7.</del> 13	I. 5	4 60 C	
7	160	0	±, ⇔ ,	& 2 24	5.0	च <del>४</del>	100 92.0 1530
9	001	2.0	500	2.5	לט ה	4, 9	
မှ	100	7.0	20	60 7 E	1.5	 → v.	
**	196	7.0	* 0	4 6; c3	2.5	च च	
63	100	() ()	φ,	4 %	1.5	₹ €	
2	001		60 0	40	დ -:	44 6	380 7.5
-	100	7 0	ć	7. 5	1.5	77	170 13.5
<b>配合内容 侧合成</b>	天 然 ゴ ムスチントンチジェン共働合体ゴム	よっプクジェンゴム カーボンブッシッ・1	¥5.	は 多 権	N-オキンジュチャンスンブチャゾーシスタンカンテッド	ない リング・アンド・マー・アン・ロー・アン・ロー・アン・アン・アン・アン・アン・アン・アン・アン・アン・アン・アン・アン・アン・	部 祭 被严辱の申び、 % 20%モジュラス、 13/6 地西 路 資務、 13/6

. IA82幅/8,DBP102碱/1008 . フエノール100重盘部K対してカシュー簡40真最部や変性したノボラック型クエノール勧脂

タイヤ記号	A	В	С	D	E	F
タイヤ構造	第2図	第1図	第1図	第1図	第1図	第1図
ビードフイラーに使用したゴム配合な	<i>1</i> 61	Mi	16.3	16.5	16.7	16.8
<u>結 果</u> 高速走行性 ㎞/b 分	185	169	177	185	193	193
積 剛 性 指 数	100	7 5	93	103	112	110
乗心地性 当り指数 減衰指数	100	115	112	109	105	196
特殊耐久ドラム試験(A条件)	3 0,6 0 0 km 完走	11,000	21,500 km	30,000 ian 完定	16.500 km	3 0,0 0 0 /m 完走

第2表から明らかなように、本発明のビードフ イラーゴム組成物を使用すれば高速走行性、操縦 安定性および耐久性が従来タイヤAと同等若しく 20 一粘度をJIS K 6300に準じて測定し、更に はそれ以上になり、しかも無心地性が砦しく改善 されている。

# 実施例 2

前記第1表の配合ル5の組成でカーボンブラツ

クのみを第3裂に示した各種のカーボンブラツク に換えて泡練りしたゴム組成物について、ムーニ 奥越例1と同様に破断時の伸び、20%モジュラ シ、動的弾性率を求めた。次いでタイヤにおける 評価を第1図に示した構造のタイヤで実施例1と 同様に行なつた。結果を第3数に示す。

第 3 表

配合內容	11.	1 2	13	5	15	1 6	1 7
カーボンブラツク IA ***タ/3 DBP **2/1009	3 6 9 1	43 121	8 6 6 0	8 2 1 0 2	8 4 1 5 0	1 2 1 I 1 4	1 4 5 1 1 3
総 サ ムーニー粘度 破断時の伸び % 20%モジュラスkg/kg/ 動的弾性率kg/kg/		73 220 50.5 610	85 250 51.5 625	9 2 2 2 5 6 0. 5 7 5 0	1 2 5 2 0 0 4 5.0 5 9 5	1 0 3 2 1 0 5 3.0 7 1 5	132 190 425 530
タイヤ記号	G	Н	I	D		J	
結 果 高速走行性 畑/h 分	185	185	185 20	185 28		185	
<b>被剛性指数</b>	95	9 7	9 7	1 0 3		98	
特殊耐久ドラム試験 (A 柴 件)	18,000 lai	30,900 編 完差	30,000 完定 完定	30,000		3 0,0 0 0 /m 完走	

第3表から明らかな様に、カーボンブラツク IA \$40~130 mg/8 T. DBP \$130 me/ 100分以下であれば耐久性が一段と改善されて いる。尙、配含※15と※17のゴム組成物は、 未加硫時の硫動性が整しく劣り、押出成型が非常 5 実施例1と同様に行なつた。但し耐久性について に困難であつたために、タイヤとしての評価は省 略した。

# 実施例 3

第4段および第5表に示した配合内容により混 実施例1と同様に評価した。結果を第4表および 第5表に示す。倚、疲労寿命については、ASFMD 412 K準じASTM F型ダンベルにて定体各型症 労試験機を用い温度70℃、伸長率35%に固定 し振動を与え、弦断するまでの振動数を求めた。 25 いる。

次いでタイヤにおける評価を上記額も幾の配合ル 22~配合格26のゴム組成物および第5奏の配 合成27,28,30,32,34および36のゴム 組成物を使用して第1図に示した構造のタイヤで は特殊耐久ドラム試験(A条件)に加え、更に過 酷な条件、すなわちカーカスプライの折返し端部 に集中する歪エネルギーが実車通常走行時の約8 倍となる様な超過荷重、超過内圧の条件にてA条 練りした各種ゴム組成物について疲労寿命と他は 10 件と同様の特殊耐久ドラム試験(B条件)を行な つた。結果を第6表に示す。

> 第4~6表から明らかなように、無変性のフェ ノール系樹脂を変性したフェノール系樹脂を併用 すると、根果的にゴム組成物の耐久性が向上して

				狱	₹'	崧				
配合內容 配合格	×	18	1.9	2.0	2 1	2.2	2 3	24	2 2	2 6
火煞过去		100	160	100	100	001	100	001	100	100
カーボンブラック・1		10	2.8	5	7.5	7.5	2 2	20 20	2.5	89 6-
カンユー変性フェノール拗脂・2	**************************************	2 4	\$	12	Ġ					
フェノール資脂 *3			ø	1.2	8 7	24	1.8	1.2	9	
トール油液性フェノール樹脂*	-0 -0			•			9	1.2	80 FM	43
ストドコン製		7	~	~	€7	જા	<b>3</b> 3	63	2	8
田 略 冊		10	10	3.0	1.0	1.0	0.	10	10	10
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数 数	-	9	9	Ģ	Ç	φ	\$	కు	чo	φ
くみやメルレンルトレッン	A ///	* %	2,4	2.4	% **	લં	₹ 20	** **i	<b>८</b> ं	¢;
张										
被断略の伸び %	- 6	160	160	155	140	120	1 4 5	170	165	160
20%モジュラス 核	kg/col	7 2	2.0	55	8	23	ς>	න භ	6 6	0 4
學的解性操 医	Kg/cai	086	920	068	7 1 5	650	710	780	8 4 0	0 8 8
滅分 路 回	-	2×106	3×10°	5×10 <sup>6</sup>	4×10°	3×10°	6×10°	8×10°	4 × 1 00	1 × 1 0¢

IA82mg/8, DBP102mc/1008

フェノール100 置量部に対してトール油60重量部で変性したノボラツク型フェノール機能

フェノール100重量部に対してカシュー油40重量部で変性したノボラツク型フェノール樹脂

**無変性のノボラツク型フェノール樹脂** \* \*

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無

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2×10 <sup>6</sup> 4×10 <sup>6</sup>	×		4×10°	90 I ×9	5×10 <sup>6</sup>	4×10 <sup>6</sup>	4 × 1 0 <sup>6</sup>	6×106	5× 10 <sup>6</sup>	4×166
		1								

\*4 トルエン変性ノホラツク型フエノール値指\*5 エトリルゴム変性ノポラツク型フエノール樹脂

蘇

タイヤ記号	К	L	M	N	0	P
ビードフイラーに 使用したゴム配合系	A6.22	<i>1</i> €23	A6.24	1625	<i>M</i> .26	1627
高速走行性 Am/h	185	185	185	185	185	185
分	12	2 1	28	2 7	28	20
横剛性瘠数	98	101	105	106	106	101
特殊耐久ドラム試 験(A条件)	30,000㎞	30,000 km 完 走	30,000 im 完 走	30,000 km 完 走	30,000 km 完定	30.000 個 完 定
特殊耐久 ドラム試 験(B条件)	9,800 km ゴム破岩	13.000 ka 完 走	13,000 km 完 走	13,000 km 完 建	13,000 km ゴム破漿	11,000㎞ ゴム破験

タイヤ記号	Q	R	S	r	ប
ビードフィ ラ ー に 使用したゴム配合ル	<i>№</i> 28	<i>1</i> 530	<i>1</i> €32	1634	<i>1</i> €36
高速走行性 km/h	177 26	185 18	185 25	185 20	185 15
模剛性指数	9 6	103	101	102	9 9
特殊耐久ドラム試 験(A条件)	28,000 細 ゴム破膜	30,000㎞ 完 建	30,000 4元 元 元	30,000 ㎞ 完 定	30,000 km 完 走
特殊耐久 ドラム試 験(B条件)	6,700㎞ ゴム破壊	13,000 km 完 走	12,000㎞ ゴム破壊	13,000 km 完 走	11,500 km ゴム破壊

# 図面の簡単な説明

使用したタイヤの性能評価に利用したタイヤの側 方部分断面図、第2図は従来タイヤの側方部分断 面図である。

1,21……タイヤ、2,22……ベルト層、

3,23……カーカス層、3,23……カーカス 第1図は本発明のビードフィラーゴム組成物を 30 プライの折返し、4 , 24 ……ビード部、5 , 25……サイドウオール部、6,26……ビード フィラー、7,27……ビードワイヤー、28… …コード層、R……りム、R<sub>T</sub>……りムフランジ<sub>c</sub>

